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(54) Title: SIMULATION OF GENE EXPRESSION CONTROL USING CONNECTRONS, INTERFERENCE RNAS (IRNAS) AND SMALL TEMPORAL RNAS (STRNAS) IN PROKARYOTIC, ARCHEA AND EUKARYOTIC GENOMES

(57) Abstract: A computer method for the determination of the interaction between transient and permanent connectrons, interference RNA and small temporal RNA.

- 1 Simulation of gene expression control using
- 2 connectrons, interference RNAs (iRNAs) and
- 3 small temporal RNAs (stRNAs) in
- 4 prokaryotic, archea and eukaryotic genomes

5

- 6 Reference to Related Application
- 7 The present application is the subject of Provisional
- 8 Application Serial No. 60/347,295 filed January 14, 2002
- 9 The present application is a continuation in part of US Patent
- 10 Application Serial Number 09/866,925 filed May 30, 2001 entitled
- 11 ALGORITHMIC DETERMINATION OF FLANKING DNA SEQUENCES THAT CONTROL
- 12 THE EXPRESSION OF SETS OF GENES IN PROKARYOTIC, ARCHEA AND
- 13 EUKARYOTIC GENOMES, incorporated herein by reference.

14

- 15 The present application is an continuation in part of US Patent
- 16 Application Serial Number 10/227,568 filed August 26, 2002
- 17 entitled Determination of flanking DNA sequences that control
- 18 the expression of sets of genes in the Escherichia coli K-12
- 19 MG1655 complete genome, incorporated herein by reference.

20

21 Introduction

- 22 The connectron structure of a genome determines sets of four DNA
- 23 sequences (called C1, C2, T1 and T2) of minimum length of 15-
- 24 bases (C1 and C2 which are in the 3'UTR of a gene, T1 which is
- 25 on the 5'-side and T2 which is on the 3'-side of a set of
- 26 genes). Typical genomes have from hundreds to tens of thousands
- 27 of these tetradic relationships spread throughout the genome.
- 28 When a gene is transcribed into RNA the C1 and C2 sequences in .
- 29 the 3'UTR find the cognate T1 and T2 double-stranded DNA
- 30 sequences to form a pair of triple-stranded RNA-DNA-DNA

generalized Hoogsteen helices. The genes between T1 and T2 are 31 32 condensed into 30nm chromatin structure and they are no longer 33 open to promotion and transcription. The lifetime of each 34 connectron is proportional to the length of the shorter of the 35 two generalized Hoogsteen helices. Within a set of genes that 36 have been removed from promotability by the formation of a 37 connectron there may be genes that themselves have the same or 38 different C1/C2 sequences in their 3'UTRs. This inclusion 39 process induces a temporal dynamic because genes that are 40 included in a connectron can no longer produce the source C1/C2 41 RNA sequences to form other connectrons. One of the most 42 obvious instances of this temporal dynamic are the so-called 43 "one-shot" connectrons in which the transcription of a gene 44 produces a C1/C2 sequence pair that forms a connectron that 45 includes the transcribed gene itself thus turning off the 46 further expression of the gene. In general, however, the 47 connectron sources (i.e. the C1/C2 sequences) and the connectron 48 flanking targets (i.e. T1 and the T2 sequences) are in different 49 portions of the genome. The evolutionary configuration of each 50 genome alone determines whether the genes turned off by one 51 connectron are associated with other connectrons.

- 52 The C1/C2 sequences that are the sources of connectrons can also
- 53 bind to the DNA double-stranded sequences of other equivalent
- 54 C1/C2 sequences in the 3'UTR of other genes. Where these trip-
- 55 stranded RNA-DNA-DNA generalized Hoogsteen helices form, the
- 56 translation of the DNA into RNA is halted and no additional
- 57 C1/C2 connectron source sequences are produced. This
- 58 interference RNA (iRNA) produces an additional temporal dynamic.
- 59 Once again the lifetime of this iRNA is varies directly with the
- 60 length of the Cl and C2 sequences. Only the relative lengths of
- 61 the lifetimes distinguish iRNAs from small temporal RNAs
- 62 (stRNAs). The iRNA and stRNA modulate the temporal behavior of
- 63 the connectrons.

64 The third type sequence-determined component that produces a 65 temporal dynamic is the permanent connectron. If all the C1/C2 sources of a given connectron can be turned off by the action of 66 other connectrons, then it is called a "transient connectron". 67 68 If, however, the generation of the C1/C2 source of a connectron is controlled only by promotion of its associated gene then the 69 70 connectron is described as being "permanent". The gene and its 3'UTR are always open to transcription and hence the C1/C2 RNA 71 72 could be continually produced. Permanent connectrons have a 73 dominant role in the temporal dynamic. Since the permanent 74 connectrons cannot be altered by any subsequent connectron, RNAi or stRNA events, they act to determine in large measure temporal 75 76 activity of the whole cell. As the documentation of genes for 77 many of the different genomes publically available on the 78 National Center for Biomedical Information (NCBI) server improves, the number of permanent connectrons detected by our 79 80 basic-methods algorithm is becoming fewer and fewer.

81 An analogy will help to make the roles of the three sequence-82 determined components clearer. A musical organ is a device with 83 three control components and one notation component. There are 84 the stops that act to connect the tone-producing pipes to the 85 keys on different keyboards. The pedals act to modulate the tones produced by individual key actions at a given time. 86 87 organ the keys on different keyboards are depressed in a variety 88 of sequences to produce the melody. The pedals are depressed in 89 a somewhat slower fashion to produce different harmonies. As a 90 composition moves from one phase to another, the organist will 91 often change the pattern of the stops. The tempo of the 92 composition is mainly determined by the rapid alteration of key 93 depressions on the different keyboards. Unlike a piano or a harpsichord where such an action produces little effect, in 94 95 organ music a given key can sometimes be held down for a relatively long time. In the same way, a pedal can be depressed 96 97 for just a short time to produce just the hint of a harmony.

- 3 -

The dynamic range of organ music, especially in an ancient

99 cathedral, produces a sense of awe in most minds. The temporal 100 behavior of cell is really very similar and just as full of awe. The connectrons interact with each other to produce most of the 101 102 rapid changes in gene expression. Sets of genes (where a set 103 can be one gene or many genes) are turned-off and, when the 104 lifetime of the connectron expires, turned-on again. Since the lifetime of a connectron is determined by the length of the 105 106 minimum intersecting sequences, some connectron lifetimes are 107 very short while others are quite long. The iRNAs and the stRNA 108 produced by gene expression also have lifetimes so they too can 109 act in short-term or long-term fashions. In the same way that the pedals act to modulate the effect of the keys, the iRNAs and 110 111 stRNAs act to modulate the temporal behavior and interaction of 112 the connectrons. The different keyboards in an organ correspond 113 to the different chromosomes in a genome. Like the stops that 114 determine the major sound forms in an organ, the permanent 115 connectrons (which are most probably driven by alarm signals 116 from outside the cell) determine the major aspects of gene 117 expression behavior. In the same way that certain patterns of 118 stops will be used for toccatas and others for fugues, we can 119 expect to find permanent connectrons associated with cell-cycle, 120 change of energy sources, and even external calls for the cell to commit suicide (i.e. apoptosis). In the organ analogy the 121 122 music (i.e. the notation component) is separate from the instrument itself. The organist can bring any piece of music to 123 124 an instrument and play it. Genomes occasionally receive DNA 125 from outside sources. Although it may be stretching the analogy 126 a bit, one might argue that the cell might "play" the new DNA to 127 see if it confers any new evolutionarily advantageous 128 properties. In the basic method patent we showed that some connectrons are controlled by promoters that do not produce an 129 130 Open Reading Frame (ORF). These ORF-less transcripts do include 131 C1/C2 sequences. As we have processed more prokaryotic, archeal and eukaryotic genomes to determine their connectron structure, 132 the number of short gene-like fragments called pseudo-genes have 133 134 increased. In the eukaryotic genomes the size of the human

135 genome (i.e. 3.5 billion bases), there is still about 90% of the

- 136 genomic DNA that is not well characterized. It may be that this
- 137 is where the "music" of the cell is stored. The utilization of
- 138 this program may be able to resolve this question.
- 139 This invention is a program method for the simulation of
- 140 cellular gene expression behavior by means of the interaction of
- 141 permanent and transient connectrons along with the iRNAs and the
- 142 stRNAs.

143

144 Prior Art

145

- 146 The Prior Art disclosed in my above identified Patent
- 147 Application is incorporated herein by reference.

148

- 149 Brief Description of the Objects of the
- 150 Invention

151 152

- 153 The object of the invention is to provide a method for using
- 154 permanent and transient connectrons and/or iRNAs and stRNAs to
- 155 show how connectrons control the expression of the genes in a
- 156 cell.

157

- 158 Description of the Drawings
- 159 The above and other objects, advantages and features of the
- 160 invention will become more apparent when considered with the
- 161 following specification and accompanying drawings wherein:

- 163 Figure 1 illustrates that (a) Complex Representation of
- 164 Connectron Formation. (b) Simplified Representation of
- 165 Connectron Formation,

166

- 167 Figure 2 illustrates that using the simplified notation (a) At
- 168 first Genes c1, c2 and c3 are free to be expressed under
- 169 ordinary promotional control. (b) Then Gene b2 begins to
- 170 express thus forming a connectron that turns off the expression
- 171 of Genes c1, c2 and c3,

172

- 173 Figure 3 illustrates that (a) Gene al begins to express thus
- 174 forming a connectron that turns off the expression of Genes bl,
- 175 b2 and b3. (b) As a result, the connectron that turned off the
- 176 expression of Genes c1, c2 and c3 is eliminated at the end of
- 177 its lifetime and then Genes c1, c2 and c3 are capable once again
- 178 of being expressed under promotional control,

179

- 180 Figure 4 illustrates that (a) If Gene c2 happens to express, it
- 181 will generate a connectron that controls the expression of Genes
- 182 al and a2. (b) As a result, the connectron formed by Gene al is
- 183 allowed to expire at the end of its lifetime thus making it
- 184 possible for Genes b1, b2 and b3 to be expressed under ordinary
- 185 promotional control,

186

- 187 Figure 5 illustrates that if Gene d1 is only under promotional
- 188 control, then it will generate a permanent connectron that
- 189 controls the expression of Genes c1, c2 and c3. The permanent
- 190 connectron generated by Gene d1 will break the cycle of gene
- 191 expression control among the "a", "b" and "c" genes,

- 193 Figure 6 illustrates that (a) Gene al can exert control over
- 194 Cycle 1 while Gene b1 can exert control over Cycles 2 and 3.
- 195 (b) A portion of the C1/C2 of Gene b1 is different from the
- 196 C1/C2 of Gene al. When Gene bl expresses, the iRNA suppresses
- 197 the expression of Gene al thus modulating its control over Cycle
- 198 1,

199

Figure 7 illustrates that (a) Two connectrons that are not in conflict. (b) Gene bl cannot form a connectron. (c) Gene al can form a connectron because it includes the smaller Gene bl connectron. This is the "Paper covers rock" rule,

204

- 205 Figure 8 illustrates that (a) Gene bl cannot form a connectron.
- 206 (b) Gene bl cannot form a connectron. (c) Gene bl can form a
- 207 connectron as long as the T2 sequence of the Gene al connectron
- 208 is separated from the T1 sequence of the Gene b1 connectron,

209

- 210 Figure 9 to 13 details the structure of the computer program
- 211 that simulates the interaction of connectrons and iRNA,

212

- 213 Figure 14 is a simulation of *E. coli* using random initial
- 214 conditions, and

215

- 216 Figure 15 is a plot of the number of changes in connectron
- 217 activity during a simulation of E. coli using random initial
- 218 conditions.

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220

Description of the Invention

- 223 The interaction of the connectrons and the iRNAs and stRNAs in
- 224 the genome of a cell generates a temporal dynamic. Figure 1a
- 225 shows the complex representation of the formation of a
- 226 connectron. This representation names the chromosome on which
- 227 the control gene and the C1/C2 sequences reside, as well as
- 228 naming the chromosome on which the T1 and T2 sequences and the
- 229 target genes reside. The simplified representation in figure 1b
- 230 just shows that the control gene causes the formation of a
- 231 connectron around the target genes. Figures 2, 3 and 4 describe
- 232 the gene expression control behavior among three sets of genes -
- 233 called a, b and c. In figure 2a, at first Genes c1, c2 and c3

234 are free to be expressed under ordinary promotional control. In 235 figure 2b, Gene b2 begins to express thus forming a connectron 236 that turns off the expression for Genes c1, c2 and c3. 237 figure 3a, Gene al begins to express thus forming a connectron 238 that turns off the expression of Genes b1, b2 and b3. 239 result of this connectron formation is shown in figure 3b. 240 result of the Gene b2 being turned off the connectron that 241 turned off the expression of Genes c1, c2 and c3 is eliminated 242 at the end of its lifetime because no more RNA is being 243 generated by the expression of the Gene b2. When this 244 connectron is allowed to expire, then Genes c1, c2 and c3 are 245 capable of being expressed under ordinary promotional control. 246 Now for the sake of this example, let us consider that the newly 247 expressible Gene c2 forms a connectron that turns off the 248 expression of the Genes al and a2. This action is shown in 249 figure 4a. As a result of turning off the Genes al and a2, the 250 connectron formed by Gene al that controls the expression of 251 Genes b1, b2 and b3 is allowed to expire at the end of its 252 lifetime. In this example we have a temporal cycle of gene 253 expression control. A "b" gene turns off the "c" genes. An "a" 254 gene turns off the expression of the "b" genes. A "c" gene 255 turns off the expression of the "a" genes, etc. Once started, 256 this cycle can continue indefinitely. If one of the controlling 257 genes in this cycle is not expressed because of promotional 258 control in the cellular environment, then the cycle of gene 259 expression control will die away.

260

261 Figure 5 shows how a permanent connectron can influence the 262 behavior of the cycle shown in figures 2 through 4. 263 expression of Gene d1 is only due to events in the cellular 264 environment - not to any other connectron control. When Gene d1 265 expresses, it generates a connectron that turns of Genes c1, c2 266 and c3. With the "c" genes permanently turned off, they cannot 267 be turned off by the expression of Gene b2. Likewise because 268 the "c" genes are turned off permanently by Gene d1, the Gene c2 cannot turn off the "a" genes. In this example, the effect of 269

270 the expression of the permanent connectron is to shut off the 271 cycle of gene expression control among the "a", "b" and "c" 272 genes.

273

274 These examples are VERY simple. Real genomes are much, much 275 more complex. Typical prokaryotic, Archeal and eukaryotic genomes have from 100 to 100,000 connectrons. 276 The utility of 277 the computer method described in this patent application is that 278 it provides an experimental basis for investigating connectron-279 controlled behavior in naturally occurring and synthetic 280 conditions.

281

282

In figure 6 the cycle of gene expression control described in figures 2 to 4 is further simplified. The numbers of genes 283 284 within a connectron as well as their names have been eliminated. 285 The three-stage cycle of temporal control is now just an 286 abstract pattern. There could, of course, be more stages in the 287 cycle. For the purpose of this example, in figure 6a Gene al 288 can exert control over Cycle 1 and Gene b1 can exert control 289 over Cycles 2 and 3. For the purpose of this example, let us assume that the C1/C2 of Gene b1 is contains a portion of the 290 291 C1/C2 of Gene al, but that Gene b1 also has a unique portion of 292 its C1/C2 that controls Cycles 2 and 3. If Gene a1 expresses 293 first then it just exerts control over Cycle 1 but if Gene b1 294 expresses first then it exerts control over cycles 2 and 3. 295 addition because there is common C1/C2 sequence between Genes b1 296 and al then the iRNA of Gene bl will block the expression of the 297 C1/C2 of Gene al. In this way Gene b1 can block the control of 298 Cycle 1 by Gene al. This is a typical way in which iRNA and 299 stRNAs exert control over cellular behavior.

300

301 The interactions of the connectrons in a genome form an abstract 302 state machine. The state of the machine is determined by the 303 pattern of gene groups that are turned off. An important 304 component of this program invention is the development of a graphic capable of representing the complexity of each state as 305

well as presenting a large number of states for visual examination. In figure 13 such a graphic is presented.

308

The key element in the computer program shown in figures 9 to 12 309 310 is the set of rules for how connectrons interact. 311 shows that two connectrons that do not share any sequence 312 elements can both form. This is particularly true if the two connectrons are on different chromosomes. In figure 7b the 313 connectron generated by Gene al forms first. When Gene bl 314 expresses, its C1/C2 RNA cannot form a connectron because the 315 corresponding T1-T2 is inaccessible. Figure 7c shows that 316 although Gene b1 has formed a connectron, the connectron 317 produced by the expression of Gene al can also form. There is a 318 children's game called "Paper, Scissor, Rock". In this game 319 320 "Paper covers Rock", "Scissor cuts Paper" and "Rock breaks 321 Scissor". The application of this rule may be subjective but the physical implementation in DNA is plausible. 322 Further computational experimentation may resolve the utility of the 323 "Paper covers Rock" rule. Figure 8a shows that Gene al has 324 formed a connectron first. Therefore the later expression of 325 Gene b1 cannot form a connectron. Figure 8b shows that the 326 expression of Gene al has formed a connectron. The Cl produced 327 by the expression of Gene b1 tries to use a portion of the T2 of 328 the Gene al connectron. This type of connectron does not have a 329 plausible physical implementation. In figure 8c the two 330 connectrons share a common T2-T1 sequence. In this case the two 331 connectrons can form because there is a plausible physical 332 333 implementation - although only just.

334

Figures 9 through 13 detail the structure of the program that
simulates the interaction of connectrons. Figure 9 is the
general structure of the computer, the program, the data files
and the printing operation. Figure 10 shows the process flow of
the program. Figures 11 and 12 describe the dominant
calculation process in the program. In conjunction with the
connectron conflict resolution rules described above, this

process does the basic simulation of connectron and iRNA interaction. Along with knowledge of our basic methods patent application, someone skilled in the art should be able to take this diagram and reproduce the cell simulation behavior. Figure 13 describes the peripheral processes for generating, printing and plotting the cell simulation data that are shown in figures 14 and 15.

349 350 ·

Figure 14 shows a simulation of the E. coli genome. vertical line is one group of genes. The presence of a vertical 351 352 line indicates that the group of genes is turned off by some connectron. The horizontal lines at the right of the figure 353 show the percentage of the gene groups turned off. The lower 354 limit (i.e. the leftmost edge) of this graph is 50% of the gene 355 356 groups turned off. The two other vertical lines are 60% and 70% 357 of the gene groups turned off. Running down the page, this side-graph shows that as the simulation proceeds, between 75% 358 359 and 85% of the gene groups are turned off. The vertical stripes 360 on the left side of this graph show that the gene groups that 361 are turned off change quite rapidly and dramatically. 362 1,000 simulation states down the whole page. For the first 100 363 simulation states the lifetimes of the connectrons are 364 randomized and kept small. This corresponds to a heating phase. 365 From simulation states 101 to 200 the lifetimes of the connectrons are increased from zero to a value determined by the 366 length of the shortest match between the (C1 and T1) sequences 367 368 and the (C2 and T2) sequences. From simulation state 201 to 1,000 the simulation runs in it normal mode. The simulation 369 370 produces extraordinarily complex behavior. Part of the utility 371 of this invention is that it will enable us to study small and large, as well as simple and complex genomic systems (i.e. 372 373 cells). By varying the lifetimes of the connectrons as well as 374 the iRNA and stRNAs, it will be possible to produce a large 375 variety of behaviors.

377 Figure 15 Shows the results of doing a larger scale simulation. 378 The upper curve is the number of connectrons going into and out 379 of existence during a 1,000 state period. The lower curve is 380 rate of change in a 1,000 state period. The cellular simulation program described in this invention is relatively inexpensive to 381 382 run in terms of computer time. As a basic cellular simulation 383 tool, this invention will become a workhorse for computational 384 experimentation. The rules of interaction between the 385 connectrons, as well as the time constants associated with the 386 various connectrons and iRNAs can be easily changed.

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This invention utilizes the capabilities in application serial no. ______ filed contemporaneously herewith and entitled "Determination of interference RNAs (iRNAs) and small temporal RNAs (stRNAs) and their interaction with connectrons in prokaryotic, archea and eukaryotic genomes". The iRNAs and stRNAs play a vital role in determining the simulation of cellular dynamics.

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398 This invention shows that the ideas of connectrons and 399 interference RNA are very powerful. The computation process 400 described in our basic methods patent application generates for 401 a given genomes a number of connectrons. At first one might 402 assume that these connectrons are static entities. 403 invention demonstrates that connectrons and iRNA do indeed 404 interact with each other in a parallel yet sequential manner. 405 If a connectron once formed stayed in existence forever, then 406 there would be no temporal dynamic. It is precisely because the 407 connectrons and the iRNA constructs have (triple-stranded 408 generalized Hoogsteen helix determined) lifetimes that the whole 409 genome can exhibit responsive and regulatory behavior. Nature 410 seems to have used a large number of very simple relationships 411 (i.e. the expression of one gene turns off the expression of other genes) to produce very complex behavior. Figure 15 shows 412

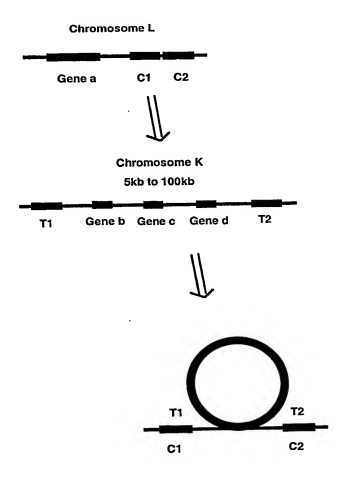
that the behavior of the *E. coli* genome is indeed very complex.

The utility of this invention will hopefully be that many
scientists throughout the world can use this tool to understand
and explore the regulatory behavior of many different genomes
ranging from the simplest bacteria through the ubiquitous (in
the sea) Archea to the plants, animals and mammals that form our
global biology.

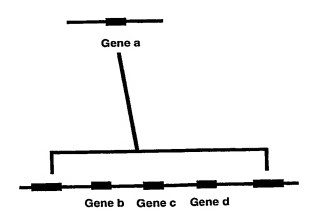
Claims

What is claimed is:

- 1. A method for using permanent and transient connectrons and/or iRNAs and stRNAs to control the expression of the genes in a cell comprising determining, by computer, the interaction of said permanent and transient connectrons and/or iRNAs and stRNAs.
- 2. A method for using permanent and transient connectrons and/or iRNAs and stRNAs to elucidate the control of the expression of the genes in a cell comprising determining, by computer, the interaction of said permanent and transient connectrons and/or iRNAs and stRNAs.

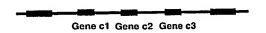


(a) Complex Representation of Connectron Formation

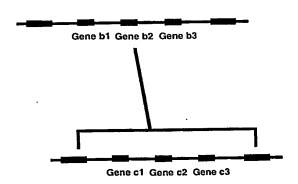


(b) Simplified Representation of Connectron Formation

Figure 1

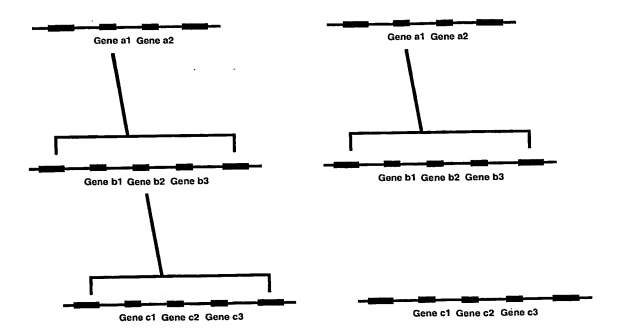


(a) At first Genes c1, c2 and c3 are free to be expressed under ordinary promotional control



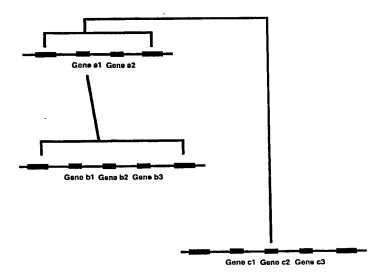
(b) Then Gene b2 begins to express forming a connectron that turns off the expression of Genes c1, c2 and c3

Figure 2



- (a) Then gene a1 begins to express forming a connectron that turns off the expression of Genes b1, b2 and b3
- (b) As a result the connectron that turned off the expression of Genes c1, c2 and c2 is eliminated at the end of its lifetime and then Genes c1, c2 and c3 are capable once again of being expressed under promotional control

Figure 3



(a) If Gene c2 happens to express, it will generate a connectron that controls the expression of Genes a1 and a2

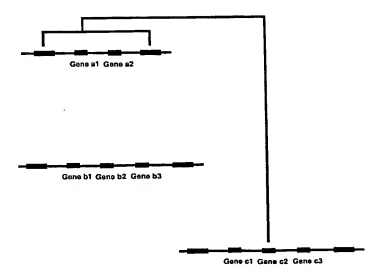
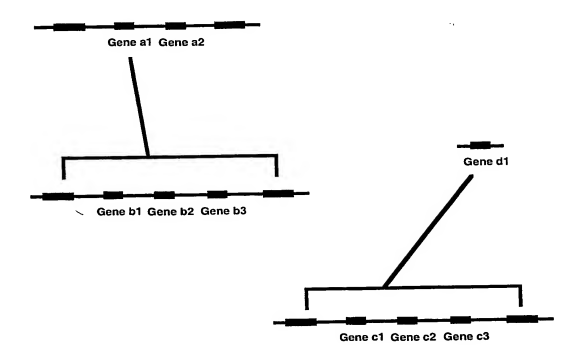


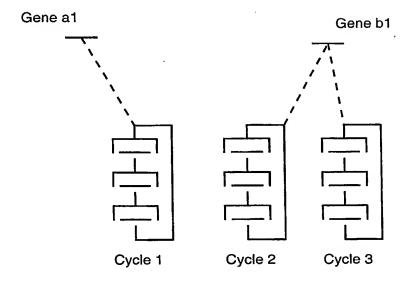
Figure 4

(a) As a result, the connectron formed by Gene a1 is allowed to expire at the end of its lifetime thus making it possible for Genes b1, b2 and b3 to be expressed under ordinary promotional control.



If Gene d1 is only under promotional control, then it will generate a permanent connectron that controls the expression of Genes c1, c2 and c3. The permanent connectron generated by Gene d1 will break the cycle of gene expression control among the a, b and c genes.

Figure 5



(a) Gene a1 can exert control over Cycle 1 while Gene b1 can exert control over Cycles 2 and 3.

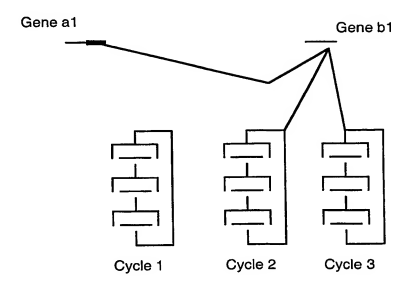
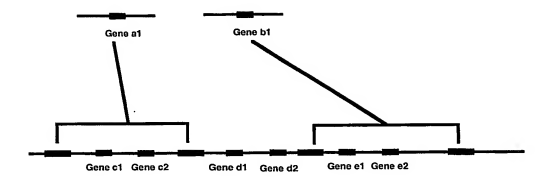
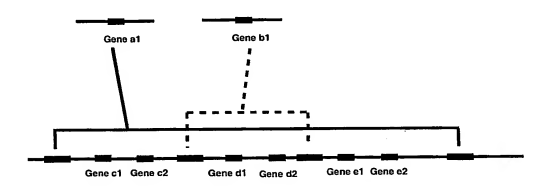


Figure 6

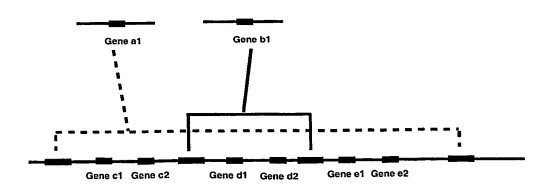
(b) A portion of the C1/C2 of Gene b1 is different from the C1/C1 of Gene a1. When Gene b1 expresses, the iRNA suppresses the expression of Gene a1 thus modulating its control of Cycle 1.



(a) Two connectron that are not in conflict

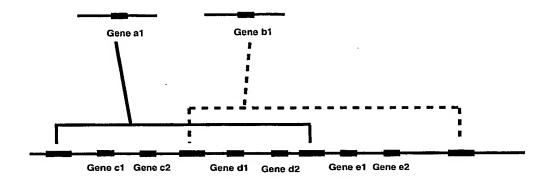


(b) Gene b1 cannot form a connectron

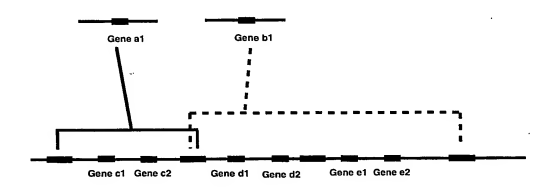


(c) Gene a1 can form a connectron because it includes the smaller Gene b1 connectron. This is the "Paper covers Rock" rule.

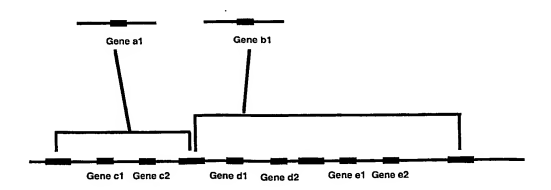
Figure 7



(a) Gene b1 cannot form a connectron



(b) Gene b1 cannot form a connectron



(c) Gene b1 can form a connectron

Figure 8

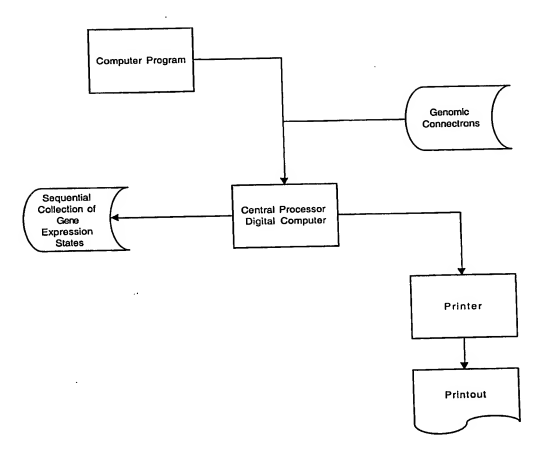


Figure 9

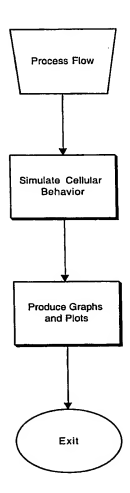
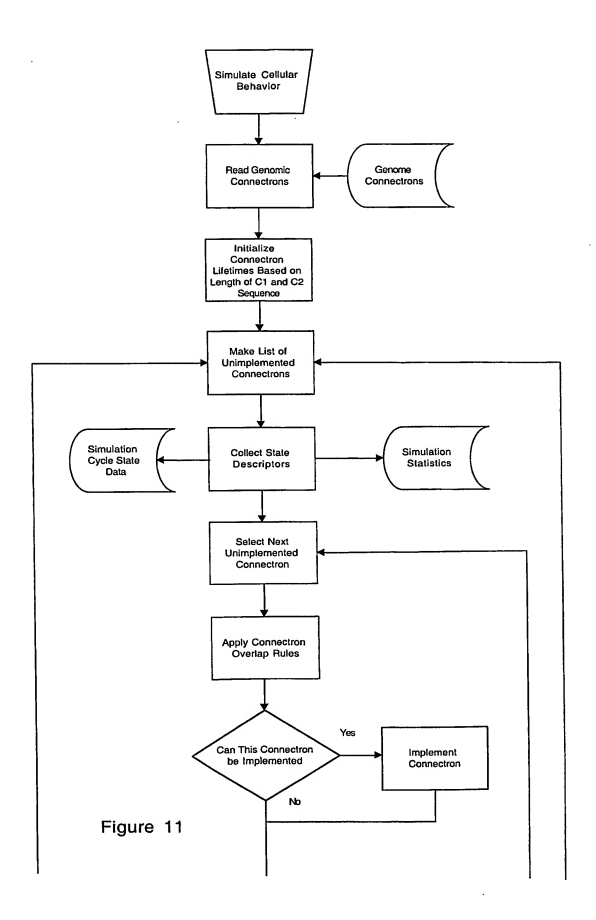


Figure 10



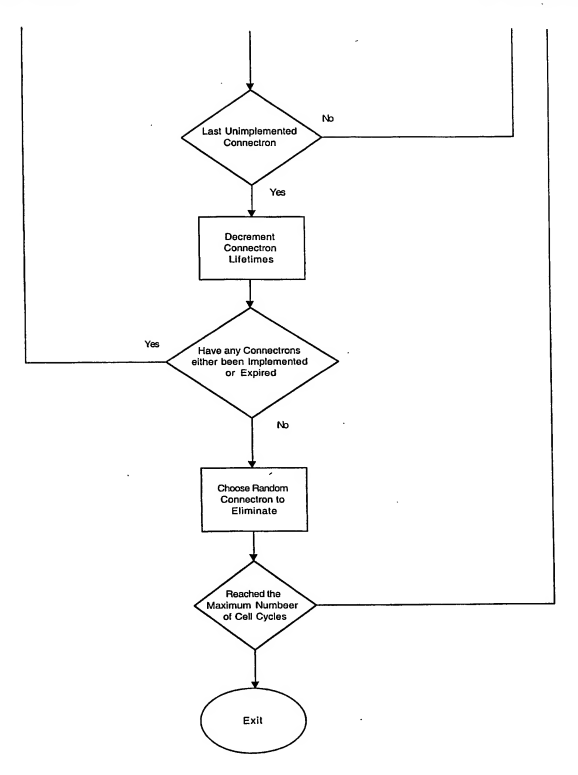


Figure 12

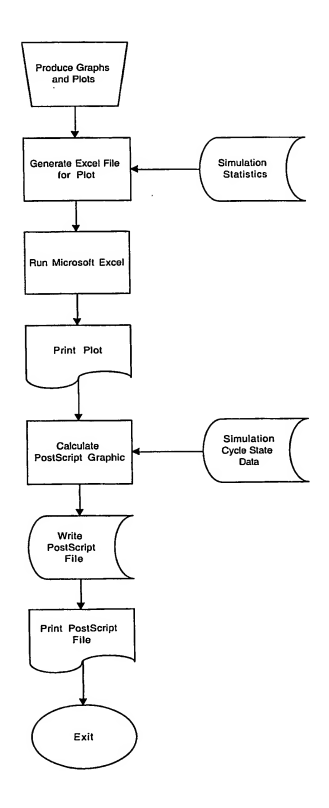
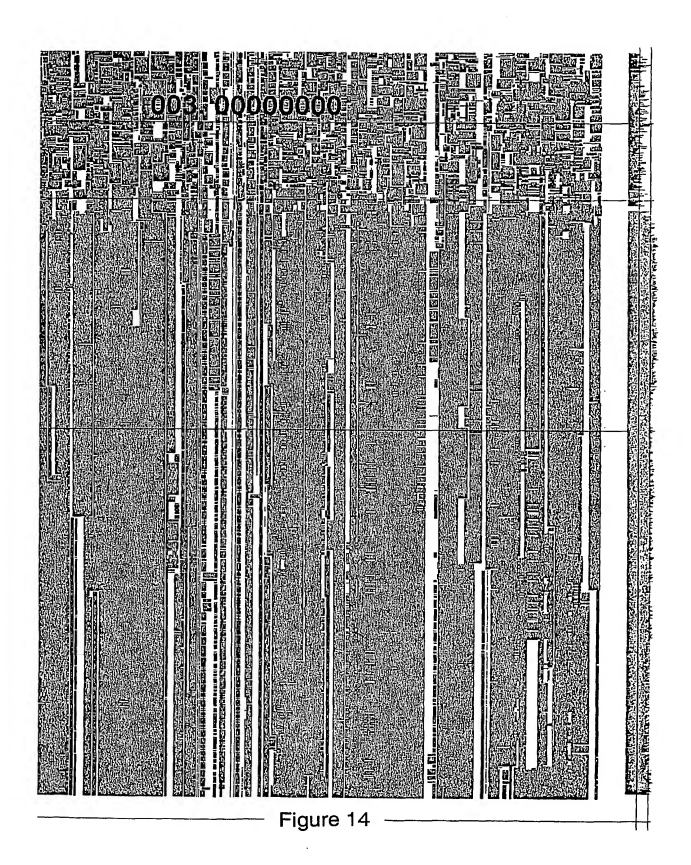
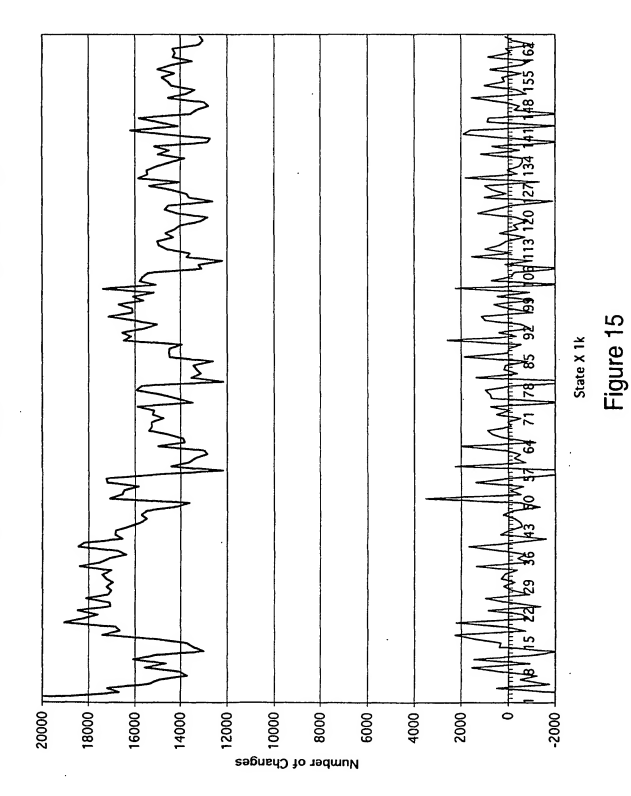


Figure 13



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E. coli - 1m Simulation - 07/05/02 - afternoon - Tay



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A. CLASSIFICATION OF SUBJECT MATTER					
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C. DOCUMENTS CONSIDERED TO BE RELEVANT					
Category *	Citation of document, with indication, where ap		Relevant to claim No.		
A	MATTICK, J.S. Non-coding RNAs: the architects of Reports, 2001, Vol. 2, No. 11, pages 986-991, espe		1, 2		
x	WO 01/094542 A2(GLOBAL DETERMINANTS, INC.) 13 December 2001		1, 2		
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